

# (12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization

# 

PCT (43) International Publication Date 11 October 2001 (11.10.2001)

(10) International Publication Number WO 01/74358 A1 60030 (US). LEE, Dennis, V. [US/US]; 2560 Highmoor

Road, Highland Park, IL 60035 (US).

9/72, 9/14, 9/20, 9/48, A61P 15/12

(51) International Patent Classification?: A61K 31/4375,

(74) Agents: KATZ, Martin, L. et al.; Rockey, Milnamow & Katz, Ltd., 47th Fibor, Two Prudential Plaza, Chicago, II. PCT/US01/40294 (21) International Application Number:

(22) International Filing Date: 14 March 2001 (14.03.2001)

60601 (US).

(8)

English

(25) Filing Language:

(26) Publication Language:

20 March 2000 (20.03.2000) US (30) Priority Data: 60/190,540 Ē (71) Applicant ffor all designated States except US): TAP HOLDINGS, INC. [115/US]; 675 North Field Drive, Lake Forest, H. 60045 (US).

Inventors; and (2) 

US/USI; 33974 North Lake Shore Drive, Gages Lake, IL. K. HISUUSI, 6986 Bennington Drive, Gurnee, IL 60031 (US). BOLLINGER, John, Daniel (US/US); 423 7th Avenne, Libertyville, II. 60048 (US). CHEN, Yisheng ZHENG, Jack, Yuqun [US/US]; 29647 N. Birch Avenue, Lake Bluff, IL 60044 (US). RELLAND, Thomas, L. [USAUS]; 1220 Vista Drive, Gumee, IL, 60031 (US). laventars/Applicants (for US only): GUPTA, Pramod

81) Designated States funtionally: AU, AG, AL, AM, AT, AU, AZ, BA, BB, GB, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DB, CM, DM, CM, EB, FR, GB, GD, GB, GH, GM, HR, HU, DD, HL, NI, SI, PK ER, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SB, SG, SI, SK, SI, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW. Linglish

Designated States (regional): ARIPO patent (GH, GM, KE, I.S, MW, MZ, SD, SL, YZ, UG, ZW), Burasian patent (AT, BB, CH, CY, DB, DK, BS, H, FR, GB, GR, IB, IT, LM, MC, N1, PT, SF, TR), OAPI patent (BF, BJ, CF, CG, CL, CM, GA, GW, ML, MR, NE, SN, TD, TG). patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European

15

Published:

with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

20

emests. Methods of administration are intranasally, be inhalation to the lungs or by oral ingestion.

PCT/US01/40294 WO 01/74358

# Methods for Treating Sexual Dysfunction With Apomorphine at Specified Plasma Concentration Levels

# Field of the Invention

S

apomorphine to a patient for the treatment of sexual dysfunction while reducing undesirable side effects. In the method, the concentration of apomorphine is ittained within the patients' plasma of up to 10 nanograms per milliliter The present invention is directed to a method for administering

Advantageously, this concentration may be achieved with less than 15% of patients so treated experiencing emesis. Methods of administration are intranasally, by inhalation to the lungs or by oral ingestion 2

### Background of the Invention

multiple doses to achieve desired results, are non-invasive and allow a rapid and convenient and simple to use, do not require a constant dosage regimen or even predictable capacity for sexual function on demand and in response to normal The human sexual response in both males and females results from a influences. Efforts are ongoing to provide effective treatments which are complex interplay of psychological, hormonal and other physiological exual stinulation For males, methods involving various external devices for the treatment of inflatable, spring driven or hydraulic models, have been used for some time. 2,818,855). In addition, penile implants, such as hinged or solid rods and impotence have been suggested such as tourniquets (see U.S, Patent No.

25

Drug treatments are also known. For example, U.S. Patent No. 4,127,118 appropriate vasodilator, in particular, an adrenergic blocking agent or a smooth nitroprusside, phenoxybenzamine, or phentolamine and a carrier to assist 4,801,587 discloses the application of an ointment to relieve impotence. muscle relaxant to effect and enhance an erection, and U.S. Patent No. ointment consists of the vasodilators papaverine, hydralazine, sodium discloses a method of treating male impotence by local injection of an

8

absorption of the primary agent through the skin. U.S. Patent No. 5,256,652

WO 01/74358 PCT/US01/40294

2

discloses the use of an aqueous topical composition of a vasodilator such as papaverine together with hydroxypropyl- $\beta$ -cyclodextrin.

The effect of apomorphine on impotence, or male sexual dysfunction has been extensively studied and reported upon. However, apomorphine has been shown to have very poor oral bioavailability. See, for example, Baldessarini et al., in Gessa et al., eds., Apomorphine and Other Dópaminomimetics, Basic Pharmacology, Vol. 1, Raven Press, N.Y. (1981), pp. 219-228.

2

Therefore, the efficacy of the use of apomorphine for treatment of sexual dyskinction is reduced by the problems of low bioavailability and undesirable, sidé effects. An increased bioavailability leads to an incréase in plasma concentration of the drug and an increase in undesirable, side effects. Therefore, for the treatment of sexual dysfunction, use of apomorphine has to date been qualified by specific concentration parameters and/or niethods of administration to overcome this problem.

2

For example, apomorphine has been disclosed for the annelioration of female sexual dysfunction in U.S. Patent No. 5,945,117. Apomorphine has also been disclosed for the amelioration of male erectile dysfunction in U.S. Patent Nos. 5,624,677, s. 1888,534; 5,770,606; 5,985,889 and 5,994,363. In U.S. Patent No. 5,624,677, mint flavoring may be added to the formulation to attenuate some of the local emesis receptors. In U.S. Patent No. 5,888,534, a slow release sublingual tablet is disclosed. The slow release of the tablet is said to reduce the undesirable side effects of the drug. The adverse effects of apomorphine were minimized by gradual acclimatization to apomorphine as disclosed in U.S. Patent No. 5,994,363. Apomorphine was disclosed for treatment of impotence in a fast release oral formulation when the patient was first pre-treated with domperidone in WO 98/31368. The treatment of erectile dysfunction with certain nasal formulations of apomorphine is disclosed in WO 99/27905.

20

15

In U.S. Patent Nos. 5,770,606 and 5,985,889 sublingual administration of apomorphine such that a plasma concentration of no more than 5.5 ng/ml was maintained was disclosed to alleviate undesirable side effects. Moreover, the '889 patent indicates that though apomorphine was evaluated as an aqueous

30

WO 01/74358

PCT/US01/40294

83

intranasal spray in Pilot Study #3, one patient's highly adverse reaction led to discontinuation of further testing and a recognition that there is still a need for reliable and relatively safe dosage formulations.

Therefore, there is a need for alternative methods of administration of apomorphine which provide the requisite bioavailability, while minimizing undesirable side effects.

We have now discovered that other routes of administration may provide a higher bioavailability than the bioavailability obtained from conventional sublingual treatment and yet do not result in a proportional increase in undesirable side effects contrary to principles understood by those skilled in the

10

### Summary of the Invention

The present invention is directed to methods for administering apomorphine to a patient for the treatment of sexual dysfunctions while reducing undesirable side effects. In the methods, apomorphine is maintained at a concentration within the patients' plasma of up to 10 nanograms per milliliter. More particularly, the present invention is directed to a method of treating sexual dysfunction in a patient comprising

15

administering a therapeutically effective amount of apomorphine or a pharmaceutically acceptable salt thereof to said patient intranasally, by inhalation to the lungs or by oral ingestion; wherein a concentration of said apomorphine is attained within said patient's plasma of up to 10 nanograms per milliliter;

20

and wherein said concentration is achieved with less than 15% of patients so treated experiencing emesis.

The present invention is also directed to a method of treating sexual dysfunction in a patient comprising administering a therapeutically effective amount of apomorphine or a

25

25

pharmaceutically acceptable salt thereof to said patient;
wherein a concentration of apomorphine is attained within said patient's
plasma of up to 10 nanograms per milliliter;

WO 01/74358 PCT/US01/40294

and wherein said concentration is achieved with less than 15% of patients so treated experiencing emesis;

with the proviso that administration is not sublingual.

The apomorphine may be administered intranasally, by inhalation to the

lungs, or by oral ingestion.

5

Intranasal administration may be accomplished by the use of a nasal spray, nasal drops, gel, suspension, ointment, cream or powder.

"Ingested orally" or "oral ingestion" as used herein indicate that the drug will primarily be pushed beyond the mouth to the stomach; so that the mouth is the point of entry but not the primary point of absorption. Thus, the terms "ingested orally" or "oral ingestion" as used herein are meant to distinguish a primarily oral absorption wherein the mouth is the point of entry and absorption occurs primarily in the stomach, from oral-mucosal administration wherein the mouth is both the point of entry and the point of absorption, or oral administration of fast dissolving tablets wherein the mouth is the point of entry but the mouth and mucosal membranes are the point of absorption. The apomorphine may be orally ingested in the form of a solution, suspension, drops, a gel, a tablet, granules, sprinkles, pills, powder or a capsule.

15

2

For the practice of any of the methods of this invention, the sexual dysfunction may be erectile dysfunction. The concentration may be attained without substantial adverse effects, such as emesis. Specifically, the concentration may be achieved with less than 15% of patients so treated experiencing emesis. The method for treating sexual dysfunction may be utilized to treat either males or females. For the practice of any of the methods of the present invention, the plasma concentration of apomorphine may preferably be from about 0.1 to about 7 ng/ml. Presently most preferably, the plasma concentration of apomorphine may be from about 0.5 to about 5 ng/ml.

25

20

The present invention is also directed to a method of treating sexual dysfunction in a patient comprising

spomorphine or a pharmaceutically acceptable salt thereof to said

intranasally administering a therapeutically effective amount of

39

PCT/US01/40294

WO 01/74358

natient

wherein a concentration of apomorphine is attained within said patient's plasma of up to 10 nanograms per milliliter.

For the intranasal route, the apomorphine may be administered as a nasal

spray, nasal drops, gel, suspension, ointment, cream or powder.

The present invention is also directed to a method of treating sexual dysfunction in a patient comprising

administering a therapeutically effective amount of apomorphine or a pharmaceutically acceptable salt thereof to said patient by oral

ingestion;

2

wherein a concentration of apomorphine is attained within said patient's plasma of up to 10 nanograms per milliliter.

For oral ingestion, the anomorphine may be administered as a solution, a suspension, drops, a gel, a tablet, pills, powder, granules, sprinkles or a capsule.

The present invention is also directed to a method of treating sexual dysfunction in a patient comprising

15

administering a therapeutically effective amount of apomorphine or a plarmaceutically acceptable salt thereof by inhalation to the lungs of said patient;

wherein a concentration of apomorphine is attained within said patient's plasma of up to 10 nanograms per milliliter.

Ŕ

The delivery device or the method of administration for inhalation may include metered dose inhalers, dry powder inhalers, nebulization of a solution or suspension and/or any other system which achieves the same results.

Detailed Description of the Invention

25

In males, the form of sexual dysfunction is erectile dysfunction. A normal erection occurs as a result of a coordinated vascular event in the penis. This is usually triggered neurally and consists of vasodilation and smooth

muscle relaxation in the penis and its supplying arterial vessels. Arterial inflow causes enlargement of the substance of the corpora cavernosa. Venous outflow

PCT/US01/40294 WO 01/74358

by reflex mechanisms. Erectile mechanics are substantially similar in the female penis sufficient to cause rigidity. Muscles in the perineum also assist in creating is trapped by this enlargement, permitting sustained high blood pressures in the nervous system by sexual thoughts or fantasy, and is usually reinforced locally and maintaining penile rigidity. Erection may be induced centrally in the

physiological abnormalities in general (organic), from neurological disturbances foregoing. Impotence may be hormonal, congenital, vascular or partial ability, neurogenic), hormonal deficiencies (endocrine) or from a combination of the Impotence or male erectile dysfunction is defined as the inability to achieve and sustain an erection sufficient for intercourse. Impotence in any given case can result from psychological disturbances (psychogenic), from among others.

2

impotence is defined as functional impotence with no apparent overwhelming spontaneous early morning, video erotica, etc.) but not others (e.g., partner or standardized method of diagnosis or treatment. As used herein, psychogenic organic basis. It may be characterized by an inability to have an erection in These descriptions are not exact, however. There is currently no response to some stinuli (e.g., masturbation, spontaneous nocturnal, spousal attention).

20

15

Females also can have sexual dysfunction that increases with age and is arterial inflow which engorges the vagina and increases vaginal Jubrication and genital response. It is known that in women, sexual arousal is accompanied by erection in the male are believed to be similar vasculogenic factors in female associated with the presence of vascular risk factors and onset of menopause. Some of the vascular and muscular mechanisms that contribute to penile that the muscles in the perineum assist in achieving clitoral erection.

25

dysfunction includes a failure to attain or maintain vaginal lubrication-swelling esponses of sexual excitement until completion of the sexual activity. Organic psychogenic causes or from a combination of the foregoing. Female sexual In the female, sexual dysfunction can arise from organic and

3

PCT/US01/40294 WO 01/74358

emale sexual dysfunction is known to be related in part to vasculogenic impairment resulting in inadequate blood flow, vaginal engorgement insufficiency and clitoral erection insufficiency.

Apomorphine ((R)-5,6,6a,7-tetralydro-6-mellyl-4II-dibenzo-[de,g]quinoline-10,11-diol) can be represented by the formula

and exists in a free base form or as an acid addition salt. For the purposes of the pharmacologically acceptable moieties forms of apomorphine can be utilized as present invention, apomorphine hydrochloride is preferred, however other

2

commensurate with a reasonable benefit/risk ratio. Pharmaceutically acceptable acceptable salt" means those salts which are, within the scope of sound medical mimals without undue toxicity, irritation, allergic response and the like and are pharmaceutically acceptable salts in detail in J. Pharmaceutical Sciences, 1977, purification of the compounds of the invention or separately by reacting a free oase function with a suitable organic acid. Representative acid addition salts Apomorphine can be used in the form of pharmaceutically acceptable salts derived from inorganic or organic acids. The phrase "pharmaceutically judgement, suitable for use in contact with the tissues of humans and lower 66. I et seq. The salts can be prepared in situ during the final isolation and salts are well-known in the art. For example, S. M. Berge et al. describe include, but are not limited to acetate, adipate, alginate, citrate, aspartate, ន 15

PCT/US01/40294

20

S

10

Apomorphine has been disclosed as useful in intranasal formulations for the treatment of Parkinson's disease in U.S. Patent No. 5,756,483.

2

Apomorphine transdermal administration has been disclosed in U.S. Patent No. 5,939,094; and apomorphine in capsule form has been disclosed in U.S. Patent

2

Apomorphine is a dopamine receptor agonist that has a recognized use as dopamine receptor agonist is administered in an amount sufficient to excite cells an emetic when administered subcutaneously in about a 5 milligram dose. For in the mid-brain region of the patient but with minimal side effects. This cell excitation is believed to be part of a cascade of stimulation that is likely to the purposes of the present invention, apomorphine or a similarly acting include neurotransmission with serotonin, dopamine and oxytocin.

25

Apomorphine according to the invention can be administered as a nasal spray, nasal drop, suspension, gel, ointment, cream or powder. The 9

PCT/US01/40294 WO 01/74358

administration of the nasal composition may also take place using a nasal tampon or nasal sponge. Powders can be administered using a nasal insufflator. Powders can also be used in such a manner that they are placed in a capsule. The capsule is set in out the powder particles. Powder formulations can also be administered in a jetan inhalation or insufflation device. A needle is penetrated through the capsule to make pores at the top and the bottom of the capsule, and air is sent to blow spray of an inert gas or suspended in liquid organic fluids.

pharmaceutically acceptable salts thereof and a physiologically tolerable diluent. salts thereof formulated into compositions together with one or more non-toxic The present invention includes apomorphine and pharmaceutically acceptable hat are collectively referred to herein as diluents, for intranasal delivery or for physiologically tolerable or acceptable diluents, carriers, adjuvants or vehicles dysfunction with a pharmaceutical composition comprising apomorphine and The present invention provides a method for the treatment of sexual oral administration in solid or liquid form.

15

also be desirable to include isotonic agents, for example sugars, sodium chloride for example, parabens, chlorobutanol, phenol, sorbic acid, and the like. It may microorganisms can be ensured by various antibacterial and antifungal agents, These compositions can also contain adjuvants such as preserving, wetting, emulsifying, and dispensing agents. Prevention of the action of

2

aluminum metahydroxide, bentonite, agar-agar and tragacanth, or mixtures of polyoxyethylene sorbitol and sorbitan esters, microcrystalline cellulose, Suspensions, in addition to the active compounds, may contain suspending agents, as for example, ethoxylated isostearyl alcohols, liese substances, and the like.

25

Among the pharmaceutically acceptable surfactants are polyoxyethylene castor polyoxyl 35 castor oil (CREMOPHOR EL), or poloxyl 40 hydrogenated castor oil derivatives, such as polyoxyethylene-glycerol-triricinoleate, also known as Useful intranasal formulations contain a stabilizer and a surfactant.

PCT/US01/40294

S

2

Among the pharmaceutically useful stabilizers are antioxidants such as sodium sulfite, sodium metabisulfite, sodium thiosulfate, sodium formaldehyde sulfoxylate, sulfur dioxide, ascorbic acid, isoascorbic acid, thioglycerol, thioglycolic acid, cysteine hydrochloride, acetyl eysteine, ascorbyl palmitate, hydroquinone, propyl gallate, nordihydroguaiaretic acid, butylated hydroxytoluene, butylated thydroxyanisole, alpha-tocopherol and lecitlin.

Preferably, the stabilizer will be between about 0.01% and 5% by weight of the plarmaceutical composition.

<u>.</u>

Chelating agents such as ethylene diamine tetraacetic acid, its derivatives and salts thereof, dihydroxyethyl glycine, citric acid and tartaric acid among others may also be utilized.

20

Proper fluidity can be maintained, for example, by the use of coating materials such as lecitlin, by the maintenance of the required particle size in the case of dispersions and by the use of surfactants.

Solid dosage forms for oral administration include capsules, tablets, pills, powders and granules. In such solid dosage forms, the active compound may be mixed with at least one inert, pharmaceutically acceptable excipient or carrier, such as sodium citrate or dicalcium phosphate and/or a) fillers or extenders such as starches, lactose, sucrose, glucose, mannitol and silicic acid; b) binders such as carboxymellyIcellulose, alginates, gelatin, polyvinylpyrrolidone, sucrose and acacia; c) lumectants such as glycerol; d) disintegrating agents such as agaragar, calcium carbonate, potato or tapioca starch, alginic acid, certain silicates

25

30

PCT/US01/40294

WO 01/74358

Ξ

and sodium carbonate; e) solution retarding agents such as paraffin; f) absorption accelerators such as quaternary annmonium compounds; g) wetling agents such as cetyl alcohol and glycerol monostearate; h) absorbents such as kaolin and bentonite clay and i) lubricants such as talc, calcium stearate, magnesium stearate, solid polyethylene glycols, sodium lauryl sulfate and mixtures thereof. In the case of capsules, tablets and pills, the dosage form may also comprise buffering agents.

Š

Solid compositions of a similar type may also be employed as fillers in soft and hard-filled gelatin capsules using such excipients as factose or milk sugar as well as high molecular weight polyethylene glycols and the like.

2

The solid dosage forms of tablets, dragees, capsules, pills and granules can be prepared with coatings and shells such as enteric coatings and other coatings well-known in the pharmaceutical formulating art. They may optionally contain opacifying agents and may also be of a composition such that they release the active ingredient(s) only, or preferentially, in a certain part of the intestinal tract, optionally, in a delayed manner. Examples of embedding compositions which can be used include polymeric substances and waxes.

15

The active compounds can also be in micro-encapsulated form, if appropriate, with one or more of the above-mentioned excipients.

Liquid dosage forms for oral administration include pharmaceutically acceptable emulsions, solutions, suspensions, syrups and elixirs. In addition to the active compounds, the liquid dosage forms may contain inert diluents commonly used in the art such as, for example, water or other solvents, solubilizing agents and emulsifiers such as ellyl alcohol, isopropyl alcohol, ellyl carbonate, ethyl acetate, benzyl alcohol, benzyl benzoate, propylene glycol, 1,3-butylene glycol, dimethyl formamide, oils (in particular, cottonsecd, groundnut, corn, germ, olive, castor and sesame oils), glycerol, tetrahydrofurfuryl alcohol, polyethylene glycols and fatty acid esters of sorbitan and mixtures thereof.

Besides inert diluents, the oral compositions may also include adjuvants such as wetting agents, enulsifying and suspending agents, sweetening, flavoring and perfuming agents.

Prescott, Ed., Methods in Cell Biology, Volume XIV, Academic Press, New Methods to form liposomes are known in the art. See, for example, York, N.Y. (1976), p. 33 et seq.

9

hereinafter in the Examples. These Examples are presented to describe preferred upon the route of administration or formulation of the drug is described in detail The moderation of undesirable side effects of apomorphine depending embodiments and utilities of the invention and are not meant to limit the invention unless otherwise stated in the claims appended hereto.

13

#### Example 1

2

studied in dogs. Dogs have been shown to be an appropriate model for study as known to be 5 to 10 times more sensitive than humans to apomorphine-induced administered intranasally as compared to the conventional sublingual route was apomorphine sub-lingual tablets in dogs have been shown to be comparable to the bioavailability by the same route of administration in humans. Dogs are disclosed in U.S. Patent No. 5,994,363 Example 3. The bioavailability of The moderation of undesirable side effects of apomorphine when

25

The drug was administered intranasally by inserting drops into the noses listed in Table 1. The intrapasal dose per dog was 2 mg in a volume of 0.2 ml. of each of a group of six dogs in amounts and three different formulations as Dogs were anesthetized lightly to avoid sneezing reflex. At each of the

30

PCT/US01/40294

WO 01/74358

13

indicated in the table. For example 2/4 in the table indicates that two dogs of a obtained in a previous study, wherein a group of four dogs were monitored for emesis after the same time intervals after administration by various routes. indicated times, the animals were checked for emesis. At a given time, the group of four had emesis at a given time. This data was compared to data number of dogs having emesis out of the number of dogs in the group is stands for sublingual and SC stands for subcutaneous.

PCT/US01/40294

14

Table 1

Comparative Raw Data for Dog Emesis Subsequent to Intranasal Apomorphine Administration

9/1 × 8 Incidences of Emesis at Given Times (min) 1/6 3/6 2 2/4 ~ 2 4/4 'n 3/6 3/6 • Dose/dog (mg) 0.4 freatment SL Tablet Intranasal Intranasal Study 1 Oral. SÇ.

2

a = daia abhained from study of Example 3 of U.S. Patent No. 5,994,363.
b = Formulation of 1% dug (10 mg/nul), 3% potypropylerac/potysoxychlydrac block copolymer (PLJROMIC F127) and 1% sodium metabisulfite (stabilizer) in water c = Formulation of 1% dug (10 mg/m), 1% polycoxypropylerac/potysoxychlylera block copolymer (PLJROMIC F127) and 1% sodium metabisulfite (stabilizer) in water d = Formulation of 1% dug (10 mg/m), 15% polycoxypropylerac/polycoxychlylera block cupolymer (PLJROMIC F127), 0.6% hydroxypropyl mettyl callulose (METHOCE K100 LV, bioadhesive agent) and 1% sodium metabisulfic (stabilizer) in water c = Formulation of 0.44% drug (0.4 mg/ml) and 1% sodium metabisulfic in water

9/1

4/6

Integrated

15

9/1

4/6

9/1

Š

20

25

Table 2 below shows the analysis of the raw data provided in Table 1 above. Bioavailability is measured relative to subcutaneous administration, which provides 100% bioavailability. C<sub>max</sub> is the maximum blood plasma concentration; T<sub>max</sub> is the time from dosing until maximum blood serum concentration is obtained; average severity (AS) is calculated as total incidences of emesis over time divided by number of dogs studied, expressed as a percentage. AS/C<sub>max</sub> is a measure of severity with respect to maximal concentration of the drug. A higher AS/C<sub>max</sub> value indicates that there is a greater proportion of side effects (measured here as emesis) relative to the

39

35

WO 01/74358

15

PCT/US01/40294

amount of drug in the subject's system. Moreover, a lower AS/C<sub>max</sub> value indicates that there is a lesser proportion of side effects relative to the amount of the drug in the subject's system. Therefore, lower AS/C<sub>max</sub> values are desirable. Note also that an AS of 50% in dogs is approximately equivalent to an AS of 5%

in humans, due to the much higher sensitivity in dogs than humans.

2

Table 2 shows that the intranasal administration results in a greatly increased C<sub>max</sub> and bioavailability over sublingual administration at the same dosage level. However, contrary to conventional behavior, the increase in severity of side effects is not also proportionally increased. The last column of Table 2 illustrates this point. Therefore, intranasal administration unexpectedly results in a more effective bioavailability than sublingual administration without a proportional increase in adverse side effects.

Table 2

Analysis of Comparative Raw Data for Dog Emesis Subsequent to

# Intranasal Apomorphine Administration

S

Treatment	Dose/dog	T	C	Bioavailability	Average	AS/C <sub>max</sub>
	(mg)		(lug/gu)	(%)	Severity	
					(%)	
SC*	0.4	0.25	8.46	100	150	17.7
SL. Tabler	. 2	0.38	1.75	13.5	50	6.5
Oral*	2	0.35.	0.40	3.9	0	0
Study 1						
Intranasal 1 <sup>6</sup>	2	11.0	139.2	150.8	83	9:0
Intranasal 2°	2	0.27	161.4	126.9	83	0.5
Intranasal 34	2	0.17	1152.6	8.201	83	0.5
ລຣ	1			001	100	

 $\cong$ 

15

15

a = data obtained from study of Example 3 of U.S. Patent No. 5,994363
b = Formulation of 1% drug (III) 18% polyoxypropyletacylolyoxychiylene block
copolyner (FULRONIC F127) and 1% sodium metabisulfite (stabilizer) in water
c = Formulation of 1% drug (II 0 mg/ml.) 13% pulyoxypropyletacylolyoxychiylene block
copolymer (FULRONIC F127) and 1% sodium metabisulfite (stabilizer) in water
d = Formulation of 1% drug (II 0 mg/ml.) 13% pulyoxypropyletacylolyoxychiylene block
Examplation of 1% drug (II 0 mg/ml.) 13% pulyoxypropyletacylolyoxychiylene block
Examplation of 1% drug (II 0 mg/ml.) 15% pulyoxypropylene(Pulyolic block
K100 LV, bloadhesive agent) and 1% sodium metabisulfite (stabilizer) in water

20

e = Formulation of 0.04% drug (0.4 mg/ml) and 1% sodium metabisulfite in water

#### Example 2

25

A solution was introduced directly to the dogs' lungs through a hole made in the apomorphine is by inhalation, as compared to the conventional sublingual route. information on moderation of undesirable side effects when administration of trachea of each dog, to represent administration of an aerosolized drug which The experimental procedure of Example 1 was utilized to obtain deposits in the lungs. The results of the study are shown in Table 3.

8

1

PCT/US01/40294

#### Table 3

## Comparative Raw Data for Dog Emesis Subsequent to Apomorphine Administration by Inhalation

. ۶

			Incide	Incidences of Emesis at Given Times (min)	mesis a	Given	Times	(min)	
Treatment	Dose/dog	0	ĸ	œ	2	51	20	30	09
	(நா)								
sc	0.4			4/4		2/4			
SL Tablet	2			•	,	Z.	-	7,	]
Oral	2							ı	
Study 2				_					
Inhalation	0.5	-	4/5	,	1	ı			•
1 <sub>p</sub>						-			
Inhalation		5/5					,	,	
2°									
Inhalation	2	5/5		,		,		,	
34									_

2

a = data obtained from study of Example 3 of U.S. Patent No. 5,994,363
b = Formulation of 0.05% drug (0.5 mg/ml) and 1% sodium metabisulfite (stabilizer) in water, 1

ml per dog c = Formulation of 0.1% drug (1 mg/ml) and 1% sodium metabisulfite (stabilizer) in water; 1 ml

per dog d = Formulation of 0.2 % drug (2 mg/ml) and 1% sodium metabisulfite (stabilizer) in water; 1 ml per dog

25

2

severity of side effects is not also proportionally increased. The last column of Table 4 illustrates this point. Therefore, administration by inhalation results in proportional increase in adverse side effects. It is particularly noteworthy that bioavailability over sublingual administration at the same, as well as at lower, this method of dosage administration allows a dose proportionate increase in Table 4 below shows the analysis of the raw data provided in Table 3 dosage levels. However, contrary to conventional behavior, the increase in above. The drug administration to the lungs results in a greatly increased more effective bioavailability than sublingual administration without a

39

PCT/US01/40294

8

Cnax, an expected phenomenon, while reducing AS/Cnax, an unexpected phenomenon.

Table 4

Analysis of Comparative Raw Data for Dog Emesis Subsequent to

Apomorphine Administration by Inhalation

Trentment	Dose/dog	٦,	C II	Bioavallability	Average	AS/C
	(But)		(ng/mt)	(%)	Severity	
					(%)	
SC	0.4	0.25	8.46	100	051	17.71
SI. Tabler	2	0.38	21.75	13.5	20	6.5
Oral*	2	0.35	0,40	3.9	0	0
Study 2						
Inhalation [*	5.0	0.17	15.2	67.2	80	53
Inhalation 2*	_	0.17	31.5	62.7	100	3.2
fuhatution 3-3	2	0.17	65.1	6:19	100	2
•			_			

2

2

a = data obtained from study of Example 3 of U.S. Patent No. 5,994,363 b = Formulation of 0.05% drug (0.5 mg/ml) and 1% sodium metabisulfite (stabilizer) in water; 1 b = Formulation

20

ut per dog c = Formulation of 0,1% drug (1 mg/mt) and 1% sodium metablistifite (stabilizer) in water, 1 mt

per dog d = Formulation of 0.2 % drug. (2 mg/ml) and 1% sodium metabisulfite (stabilizer) in water, 1 ml per dog

25

Example 3

39

The experimental procedure of Example 1 was utilized to obtain

information on the moderation of undesirable side effects when apomorphine is sublingual route or oral route. Test formulations were introduced directly to the dogs' stomach as a solution through a tube or in capsule form. The results of administered orally by various formulations, as compared to the conventional the study are shown in Table 5.

WO 01/74358

19

PCT/US01/40294

Comparative Raw Data for Dog Emesis Subsequent to Table 5

Oral Apomorphine Administration

5

			무	Incidences of Emesis at Given Times (min)	Emesi	s at Giver	Тітез	(min)	
Treatment	Bop/asoQ	0	2	8	10	15	70	30	8
	(mg)								
. SC.	0.4		•	4/4		2/4			
SL Tablet	2		ı			7,	,	7/	
Oral	2		•			•			
Study 3									
Oral 1*	. 01						,	,	1/5
10mg/ml gavage									
Oral 2	20	2/2		3/5					
20mg/ml gavage									
Oral 34 copsules	10				,			,	
						•			

12

a = data obtained from study of Example 3 of U.S. Patent No. 5,994,363

15

b = Formulation of 1% drug (0.5 g) and 1% (0.5 g) sodium metabisulfite (stabilizar) in water c = Formulation of 2% drug (1 g) and 1% (0.5 g) sodium metabisulfite (stabilizar) in water d = Formulation of 10% drug (10 mg) and 90% (90 mg) of Avicel 101 (niuxocystalline celluloss)

2

25

of note is that different oral formulations produce varying Cmx values. The oral tablets without a comparable increase in entesis. Therefore, depending upon the above. The relationship of bioavailability to severity of undesirable side effects formulation 2 results in a higher bioavailability than oral formulation 1, yet oral formulation 2 produces less severe side effects in relationship to bioavailability formulation 2 resulted in nearly a four-fold higher  $C_{\max}$  compared to sublingual than oral formulation 1. The last column of Table 6 illustrates this point. Also Table 6 below shows the analysis of the raw data provided in Table 5 can be controlled by varying the formulation for oral administration. Oral

30

formulation, Cmx versus side effects can also be optimized.

PCT/US01/40294

20

# Analysis of Comparative Raw Data for Dog Emesis Subsequent to Oral Apomorphine Administration

Table 6

Treatment	Dose/dog	,i	C.	Bloavailability	Average	AS/C
	(But)		(ag/ml)	. (%)	Severity	
					(%)	
SC	0.4	0.25	8.46	001	150	7.71
SI, Tubler	2	0.38	27.7	13.5	20	6.5
Oral*	2	0.35	0.40	3.9	0	0
Study 3						
Oral 1	10	0.13	4.21	1.83	20	4.75
l'Umg/ijl gavage				٠		
Oral 2*	20	0.35	29.3	3.87	100	3.4
20mg/ml gavage						
Oral 34 capsule	01	61.0	1.75	91'1	0	8.6

9

a = data obtained from study of Example 3 of U.S. Patent No. 5,994,363

7

b= Formulation of 1% drug (0.5 g) and 1% (0.5 g) sodium metabisulfite (stabilizer) in water c= Formulation of 2% drug (1 g) and 1% (0.5 g) sodium metabisulfine (stabilizer) in water d= Formulation of 10% drug (10 mg) and 90% (90 mg) of Avicel 101 (microcystalline

23

52

varying dose levels. Twenty-four men were tested using dosages of 2, 4, 5 and 6 sublingual tablet has been shown to offer good efficacy and minimal side-effects the tongue, the remaining undissolved mass (if any) was discarded. The samples were then assayed using a highly sensitive LC/MS/MS technique. Peak plasma reported, as indicated in Table 7. In the table, SD stands for standard deviation. sampling at specified time intervals, up to 20 minutes. After 20 minutes under A study was done to determine apomorphine absorption in humans at drug levels approximating 0.70, 1.25, 1.70 and 1.91 ng/nıl respectively were These results indicate that apomorphine is absorbed in a dose-proportionate manner (Cmax as well as AUC (area under the curve) increased linearly with increase in sublingual tablet dose). Since up to a 6 mg dose delivered via a immediately after placing the tablet under the tongue, followed by further mg sublingual tablets. Plasma samples were obtained from each subject

3

WO 01/74358

PC1/US01/40294

21

apomorphine as a sublingual tablet are meaningful indicators of performance. In other words, plasma drug levels between 0 to 6 ng/ml in humans (obtained with Jysfunction. The bioavailability of sublingual tablets in humans, relative to a 6 mg tablet), following sublingual administration as a tablet, are meaningful indicators of good efficacy and low side-effects in the treatment of sexual in humans, plasma drug levels attained following administration of 6 mg subcutaneous control, was estimated to be 16-18%.

Ś

# Apomorphine Pharmacokinetic Parameters in Humans

2

Parameter		2 mg SL	4 mg SL	5 mg SL	6 mg SL	I mg SC
E	Menn	0.74	0.72	89.0	99'0	0.34
	SD	0.30	0.32	0.21	0.32	0.17
c <sub>ms</sub> (ng/ml)	Mean	0.70	1.25	1.70	161	3.22
	SS	0.37	080	1.32	173	1.67
AUC.(ng-h/ml)	Mean	1.23	137	2.92	3.60	3.39
	SD	0.48	1.06	1.50	1.73	1.09

emesis. Any formulation or dosage administration technique which allows drug levels to be attained in the range of 0.25 to 5 ng/ml with less side effects such as described. Hence, any formulation or dosage which enables drug levels in dogs intranasal, inhalation to the lungs or oral formulations investigated in this work compound in the treatment of sexual dysfunction. Dogs have been indicated to numans has demonstrated about 13% incidence of nausea and 2% incidence of emesis can be expected to improve patient compliance, and usefulness of this Clinical experience with 2 to 4 mg sublingual apomorphine tablets in comparable to that achievable with sublingual tablets without comparable emesis profile is believed to have superior performance in humans. The be much more sensitive to emesis than humans, as has been previously demonstrate that this can be achieved. 25 20 15

Sub-lingual apomorphine tablets have demonstrated approximately 15% relative bioavailability against sub-cutaneous human control in humans as well

PCT/US01/40294

22

as in dogs. This suggests that the dog is a good model in representing absorption comparable side-effects. The intranasal, inhalation to the lungs or oral routes of ng/dog were investigated to achieve plasma drug levels in dogs comparable to administration investigated in the above examples demonstrate that this can be dose in dogs. For the studies presented here, dosages in the range of  $0.5\ \text{to}\ 20$ weight, an 8 mg human dose compares well with about 1.33 mg apomorphine of apomorphine. Up to 8 mg of apomorphine tablets have been shown to be well tolerated in humans. Assuming a 60 kg human weight and a 10 kg dog or higher than those achieved with 2 mg sublingual tablets in dogs without achieved.

All references cited are hereby incorporated by reference.

2

The present invention is illustrated by way of the foregoing description illustration, since many variations will become apparent to those skilled in the art in view thereof. It is intended that all such variations within the scope and and examples. The foregoing description is intended as a non-limiting spirit of the appended claims be embraced thereby.

15

the method of the present invention described herein without departing from the Changes can be made in the composition, operation and arrangement of concept and scope of the invention as defined in the following claims:

WO 01/74358

PCT/US01/40294

23

#### Claims

We claim:

administering a therapeutically effective amount of apomorphine or a wherein a concentration of said apomorphine is attained within said 1. A method of treating sexual dysfunction in a patient comprising pharmaceutically acceptable salt thereof to said patient; patient's plasma of up to 10 nanograms per milliliter;

and wherein said concentration is achieved with less than 15% of patients so treated experiencing emesis;

with the proviso that administration is not sublingual.

2

2. The method of claim 1 wherein said apomorphine is administered intranasally. 3. The method of claim 2 wherein said apomorphine is administered as a nasal spray, nasal drops, gel, suspension, ointment, cream or powder.

15

4. The method of claim 1 wherein said apomorphine is administered by oral ingestion.

20

5. The method of claim 4 wherein said apomorphine is administered as a solution, a suspension, drops, a gel, a tablet, granules, sprinkles, pills, powder, or

6. The method of claim 1 wherein said apomorphine is administered by inhalation to the lungs.

25

7. The method of claim 6 wherein said apomorphine is administered through a metered dose inhaler, dry powder inhaler, nebulized solution or

nebulized suspension.

PCT/US01/40294

24

8. The method of claim 1 wherein said sexual dysfunction is erectile dysfunction.

9. The method of claim 1 wherein said patient is female.

3

10. The method of claim 1 wherein said concentration of apomorphine is from about 0.1 to about 7 ng/ml in said patient's plasma. 11. The method of claim 1 wherein said concentration of apomorphine is from about 0.5 to about 5 ng/ml in said patient's plasma.

2

administering a therapeutically effective amount of apomorphine or a 12. A method of treating sexual dysfunction in a patient comprising wherein a concentration of said apomorphine is attained within said intranasally, by inhalation to the lungs or by oral ingestion; and wherein said concentration is achieved with less than 15% of pharmaceutically acceptable salt thereof to said patient patient's plasma of up to 10 nanograms per milliliter; patients so treated experiencing emesis.

2

13. The method of claim 12 wherein said apomorphine is administered intranasally as a nasal spray, nasal drops, gel, suspension, ointment, cream or powder.

20

lyy oral ingestion as a solution, a suspension, drops, a gel, a tablet, pills, powder, 14. The method of claim 12 wherein said apomorphine is administered granules, sprinkles or a capsule.

25

15. The method of claim 12 wherein said apomorphine is administered by inhalation to the lungs by a metered dose inhaler, dry powder inhaler, nebulized solution or nebulized suspension.

39

WO 01/74358

PC1/US01/40294

25

16. The method of claim 12 wherein said sexual dysfunction is erectile dysfunction.

17. The method of claim 12 wherein said patient is female.

S

18. The method of claim 12 wherein said concentration of apomorphine is from about 0.1 to about 7 ng/ml in said patient's plasma. 19. The method of claim 12 wherein said concentration of apomorphine

is from about 0.5 to about 5 ng/ml in said patient's plasma. 2

INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/40294

A. CLA IPC(1) US CL	. CLASSIFICATION OF SUBJECT MALLEK IPC(1) :AGIK 31/4575, 9/79, 9/14, 9/20, 9/48; AGIP 15/19 US CL :184/ 40, 434, 435, 451, 404, 484, 489; 514/264, 944, 999		
According B. F1E	According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED	national classification and IPC	
Minimum	Minimum documentation searched (classification system followed by classification symbols)	by classification symbols)	
U.S. :	42+/ 46, 434, 435, 451, 464, 484, 489; 514/584, 944, 929	929	
Documents	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched	he extent that such documents are in	cluded in the fields
Electronic	Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)	me of data base and, where practicable	, search terms used)
WEST, STH, FI	west, uspatfull sth, file registry, index		
C. DO	DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	ropriate, of the relevant passages	Relevant to claim No.
7	US 5,945,117 A (EL-RASHIDY et al.) 31 Augus (31.08.1999), abstract, column 3, lines 28-42, lines 49-57.	et al.) 31 August 1999 s 28-42, lines 49-57.	1-19
>-	WO 99/27905 A1 (DANBIOSYST UK LIMITED) 10 June 1999 (10.06.1999), abstract, page 19, lines 10-12, claims 1, 12,13, 20-24, 26.	C LIMITED) 10 June 1999-12, claims 1, 12,13, 20-24,	1-3, 6-13, 15-19
7	US 5,472,954 A (LOFTSSON) 05 December 1995 (05.12.1995), abstract, column 18, lines 62-65, claims 1 and 21.	cember 1995 (05.12.1995), s 1 and 21.	4-5, 14
	Further documents are listed in the continuation of Box C.	See patent samily annex.	
	Special extegnites of elted documents: document deflating the general state of the art witch is not considered	"y" hater document published after the fakenestions filling date or priority date and no souther with the explication but study to meantish the principle or theory medestyling the investor.	mutional filling date or priority tiertion but elied to understand invention
la la		"X" document of particular usborance, the claimed forention acunot be considered to investive stap	e claimed forention eaunct he red to investive stap
i.	decement which may favor doubts on pelonity slatin(s) or which is eited to establish the publication date of another citation or other special resean (se specified)	when the document is taken alone "Y" document of particular relevance; the	e claimed invention annot be
þ	document referring to an oral disclosure, use, arbibition or ather motos	confidered to firely an inventive stay when the abomment is combined with one or more others area documents, each combination heling obvious to a person abilise in the art	when the document is combined nearly, each combination being
ļ.	document published prior to the international filling date but later than the priority date chained	'A" document member of the same palent family	family
Date of th	Date of the actual completion of the international search	Date of mailing of the international search report	arch report
01 JUNE 2001	E 2001	16)	16 JUL 2001
Name and Commiss Box PCT	address of the ISA/US tents and Trademarks	Authorized officer Divided	dop
Washing.	C. 20231	Telegram NGUYEN	K
Facsinitle No.	(103) 305-3330	Telephone No. (703) 508-1935	د

Bax PCT

Bax PCT

Varianglon, DC, 20231

Foriniel No. (708) 305-3230

Form PCT/ISA/210 (second sheet) [July 1998]\*